Comparison of Direct Immunofluorescence and Giemsa Staining in Chlamydia trachomatis Follicular Conjunctivitis

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1. Background

Chlamydia is a genus of bacteria that consists of intracellular obligate parasites and its members fall into one of these three species: Chlamydia trachomatis, Chlamydia psittaci or Chlamydia pneumoniae. This organism can present itself in two different forms during its life-cycle; elementary body (EB) which is the infectious form and reticulate body (RB), the non-infectious form. The EB obtains a protein component termed major outer membrane protein (MOMP) which is specific for each species and sub-species and can be identified by a monoclonal antibody (Ab). C. trachomatis is divided into 18 serotypes. Serotypes A, B, Ba and C can cause hyper-endemic blinding trachoma in humans and serotypes D, E, F, G, H, I, J and K are responsible for chlamydial pneumonia in neonates and chlamydial-related inclusion conjunctivitis, non-gonococcal urethritis, cervicitis, salpingitis, pelvic inflammatory disease, proctitis and epididymitis in infants and adults. Serotypes L1, L2 and L3 are the perpetrators of lymphogranuloma venereum [1].

Gallo Vaulet et al. conducted a study in which the distribution of C. trachomatis genotypes among symptomatic patients was evaluated. The researchers found an association between genotype E and neonatal conjunctivitis [2].

Chlamydial infections are one of the most common eye diseases in humans and can cause different types of eye infections including neonatal inclusion conjunctivitis, adult chlamydial keratoconjunctivitis and trachoma. The inflammation of conjunctival stroma in chronic conjunctivitis is presented via formation of papillae and follicles. Chronic follicular conjunctivitis or a type of conjunctivitis that lasts longer than 16 days and is associated with follicle formation in conjunctiva can be related to different etiologies which are either of infectious or non-infectious origin. Chlamydial infections are one of the infectious etiologies of chronic follicular conjunctivitis and this specific association is the area of interest in our research [3, 4].

Laboratory diagnostic techniques used for chlamydial infections are either culture-based (cell culture) or non-culture based methods. The culture method is the reference and gold standard method for chlamydia identification. The non-culture based assays for chlamydia detection are as follows: 1- Direct testing like detection of inclusion bodies via Giemsa staining and detection of chlamydia EBs by immunofluorescence staining (DFA; direct fluorescent antibody) 2- Serologic tests such as complement fixation, enzyme-
linked immunosorbent assay (ELISA), indirect immunofluorescence assay (IFA) and microimmunofluorescence (MIF). Molecular tests including polymerase chain reaction (PCR), ligase chain reaction (LCR) and DNA-RNA hybridization [1, 5].

2. Objectives

In this study we aimed to assess the prevalence of *C. trachomatis* in conjunctival specimens of patients diagnosed with follicular conjunctivitis and also compare the two direct testing techniques implemented in chlamydial conjunctivitis diagnosis, i.e. Giemsa staining and direct immunofluorescence (DIF).

3. Patients and Methods

In this descriptive and comparative study, 285 patients primarily diagnosed with follicular conjunctivitis and also clinically suspected to be infected with *C. trachomatis* were referred to the clinical laboratory of Farabi Eye Hospital for diagnosis of *C. trachomatis* infection. For all subjects in this study, the conjunctival scraping method was applied in order to obtain conjunctival specimens. For this purpose, the tarsal conjunctiva of upper and lower eyelids of both right and left eyes was scraped in a sterile fashion. Subsequently, the specimens were separately placed on regular microscopic glass slides and immunofluorescence slides and stained via the two Giemsa and DIF methods. In the Giemsa method samples were stained with Giemsa stain after air-drying and in the DIF method slides were first methanol-fixed for five minutes and then stained with a monoclonal IgG Ab conjugated with fluorescein that detects the MOMP of all 18 serotypes of *C. trachomatis* (Product name: *C. trachomatis* DFA kit; Code: PL.1010; Manufacturer: Pro-Lab Diagnostics, Canada). The Giemsa-stained and DIF-stained slides were viewed and evaluated under regular optical microscope and immunofluorescence microscope, respectively. Samples that were not sufficient enough in terms of adequate number of epithelial cells, etc. were excluded from the study and re-sampling was performed in these cases. In this study patients were not charged with any additional expense and as they referred to the lab for diagnostic purposes, routine sampling was carried out. It should be noted that no additional intervention was carried out in this study and standard sampling and staining for diagnosis of conjunctival lesions was performed upon physician request. The results of the two staining methods were recorded and the related data was statistically evaluated. In order to compare the two diagnostic methods put to use in this study, we considered the DIF assay as the standard technique in this scientific endeavor (C. trachomatis infection is confirmed when DIF is positive). Sensitivity, specificity, false positive rate and false negative rate of the Giemsa staining method were extracted. For calculating these four values, we used the following formulas:

Sensitivity = true positive/(true positive + false negative),
Specificity = true negative/(true negative + false positive),
False positive rate = 1 - specificity = false positive/(false positive + true negative),
False negative rate = 1 - sensitivity = false negative/(false negative + true positive).

### Table 1. Results of Comparison of DIF and Giemsa Staining Techniques Based on Different Age Groupsa

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*All age groups*

| Male   | 50   | 27     | 22            | 28            | 5              |
| Female | 59   | 20     | 20            | 39            | 0              |

*aAbbreviation: DIF: direct immunofluorescence.*
4. Results

A total of 285 follicular conjunctivitis patients suspected to have a chlamydial infection underwent sampling from right and left tarsal conjunctivae. A number of 109 subjects (38.24%) were positive for *C. trachomatis* when DIF was implemented (Figure 1). To be noted, 50 of them were males (45.7%) and 59 of them females (54.1%). Most of the patients who tested positive with DIF were in their third and second decades of life (21 - 30 and 11 - 20 years), with prevalence of 36.69% and 19.26% respectively.

On the other hand, a total number of 47 patients (16.49%) (27 males [57.44%] and 20 females [42.55%]) showed positive results for *C. trachomatis* when the Giemsa staining method was used (Figure 2). A more detailed depiction of the results which is based on different age groups is demonstrated in Table 1.

The false negative and false positive rates of Giemsa staining were 61.47% and 2.84%, in order. Taking DIF method into account, sensitivity and specificity of Giemsa method equaled 38.53% and 97.16%, respectively.

![Figure 1. Epithelial cells of conjunctiva containing intra-cytoplasmic inclusions labeled with fluorescent materials (yellow dots), direct immunofluorescence method, fluorescence microscope, 100× magnification.](image1)

![Figure 2. Epithelial cells of conjunctiva containing intra-cytoplasmic inclusions “draped” around nucleus, Giemsa staining method, optical microscope, 400× magnification.](image2)

5. Discussion

Among the two staining procedures used in this study, the DIF method was chosen as the standard technique; therefore, the Giemsa method faced false negative and false positive rates of 61.47% and 2.84%, respectively. In comparison to DIF, Giemsa staining’s sensitivity and specificity was 38.53% and 97.16%, in order. *C. trachomatis* is the most common sexually transmitted bacterial infection (STI) [6, 7] and its prevalence differs with age, gender, race, and residency of the infected person and type of diagnostic test [6]. Chlamydial infections are one of the most frequent ophthalmologic disorders and a study declared that among 15 - 44 year olds, 1 out of every 44,000 of them contract the infection each year and prevalence in total population was 1 in 100000 [8]. Several studies have pointed out *C. trachomatis* as the most common etiology of chronic follicular conjunctivitis and most significant cause of bacterial conjunctivitis in neonates [9]. However, the prevalence of this infection has varied from 20 - 90% in scientific literature, which is parallel to age groups and the prevalence of chlamydial eye infections positively correlates with genital disease prevalence [5, 8].

In our study most of the patients belonged to the 21 - 30 years age group and after that, 11 - 20 years age group. After the age of 50, a significant decrease in the number of positive cases was witnessed in our study and this in accord with similar studies [10-13]. Our study was mainly based on adult chronic follicular conjunctivitis and we only had one pediatric case aged less than one year because neonates usually receive treatment in child therapeutic centers and are less frequently referred to ophthalmologic hospitals. In our country, prior to our current effort, only one study has been carried out to evaluate the association between chlamydia and conjunctivitis but that assessment was limited to the neonatal age group and considered neonatal conjunctivitis. In that specific study, specimens from 170 neonates diagnosed with conjunctivitis were obtained and 10 samples (6%) tested positive for chlamydia via DIF method. The relatively low prevalence of chlamydial conjunctivitis in the neonates of that study, compared to reports from the West, was linked to a lower prevalence of STIs in Iran [10]. Regarding neonatal conjunctivitis, a study in Argentina reported a 7.8% prevalence of chlamydia-related conjunctivitis [11].

In our study from the total 285 patients referred to the lab, 109 of them (38.24%) were positive in terms of chlamydial infection. Similar studies have reported positive results of 8.6% and 38% in sample sizes of 93 and 100 patients, respectively [12, 13]. It should be noted that neither our study nor others’ detected a significant sex-based difference in chlamydial conjunctivitis contraction [2, 9, 11].

Considering the age distribution in this assessment and the fact that most of the patients were part of the more sexually active age group and even some of them had previously experienced genitourinary infections, spread of infection through genital means and by eye-hand con-
tact can be an important factor in disease transmission. Based on patients’ medical history, other factors related to disease transmission are polluted water bodies, residing in sub-urban areas, swimming pools and sharing non-hygienic cosmetic agents (especially common in young women). Various methods are available for the laboratory diagnosis of infections caused by *C. trachomatis*. These assays differ in sensitivity, specificity and positive predictive value (PPV). The most suitable test or a combination of different diagnostic evaluations is determined based on several factors: 1- the susceptible population 2- accessible facilities 3- price of consumables 4- technician’s skill 5- type of specimen [14]. Prevalence and the disease in the tested population are the most important criteria for type of test. The type of the selected sample depends on the symptoms of the disease. Generally, laboratory diagnosis of chlamydial infections is based on culture and non-culture methods.

Cell culture is the gold standard and reference method for diagnosis of chlamydial infections and must be carried out in suspicious cases. Sensitivity, specificity and PPV of the culture method vary in different reports. Based on these reports, theses have ranged 50 - 95% for sensitivity, 92 - 100% for specificity and 73 - 98% for PPV [14]. The problems associated with performing cell culture are: 1- it requires a specific transport medium, 2- cells need to be cultured by a maximum 4 - 6 hours after sampling, 4- it is a time-consuming and expensive method and 5- it is inaccessible at most labs, especially in regions where chlamydia is not endemic. Therefore, this method is currently not used as a main diagnostic technique and is especially not used when samples are recruited non-invasively [15-17]. Recent articles and reports indicate that the sensitivity of cell culture is not higher than that of other methods such as DFA, enzyme immunoassay (EIA) and nucleic acid probe (NAP)-based technique and for ophthalmologic samples; these methods are more preferred compared to other methods [15]. Another method used for diagnosis of *C. trachomatis*, especially in under-equipped centers, is the direct technique of smear preparation and staining via Giemsa method in order to view intra-cytoplasmic inclusions. This technique is especially useful in neonates and obtains a lower sensitivity for adults. On the other hand, it is easy to perform and is accessible. Ultimately, this technique has established itself as one of the primary, well-known methods in our country, even though it does not have high sensitivity.

A serologic technique is less useful in diagnosing trachoma and inclusion bodies but is the test of choice for neonatal infections and diagnosing pneumonia caused by *C. trachomatis* in children. In general, serologic methods face some limitations due to low sensitivity and specificity and should be confirmed by other assays [7, 12] because in many individuals the antibodies that are found are related to a previous infection and because chlamydial infections are mostly localized, a 4-fold increase in Ab titer is not useful for diagnosis. Also, the Ab titer may not rise until two weeks and this could cause an unacceptable delay in diagnosis of the disease. However, with advances in serologic methods and using specific responses to specific groups of *C. trachomatis* antigens via MIF method, it is possible to detect a novel infection in patients who were previously infected with another immunotype [15]. Using non-culture and non-amplified techniques such as DNA probe test is specifically reserved for situations where PCR and culture methods are not available [18]. Nucleic acid amplification tests (NAATS) e.g. PCR obtain the highest sensitivity and their specificity is approximately equal to cell culture. Due to its high sensitivity and high specificity, NAAT is more effective for non-invasive samples. However, using this technique for routine diagnostic purposes regarding eye infections is costly and the technique is mostly used in research [19, 20]. DIF technique is the most suitable method for diagnosing chlamydial conjunctivitis. This rapid test has a relatively high sensitivity (70 - 100%) compared to cell culture. The specificity of the test exceeds 95% (87 - 99%). It is the only test that allows evaluation of specimen adequacy. Samples containing 10 - 20 columnar and/or squamous metaplastic cells are acceptable. Antibodies against MOMP in chlamydia are species-specific. This technique is very suitable for routine diagnostic means [15].

Taking the aforementioned characteristics of DIF and its high sensitivity and high specificity into account, we considered the DIF method as the standard test of choice in our study and the other diagnostic test used in our study, i.e. Giemsa staining (which is the current method used in most labs) was compared to it. A few similar studies have been conducted in this area (comparing different non-culture methods, especially the two direct testing techniques, DIF and Giemsa staining) and most of them have evaluated women’s vaginal samples. In a study carried out by Palayekar et al. the two direct tests, i.e. DIF and Giemsa staining were compared to each other. In total, 16.8% and 10% of the samples were DIF-positive and Giemsa-positive, respectively [21]. In another study, the sensitivity of DIF and Giemsa method for identifying *C. trachomatis* inclusions in McCoy cells was assessed. The results showed higher sensitivity of DIF in comparison to Giemsa [22]. In a study performed by Lin et al. [13], rapid diagnostic methods for chlamydial conjunctivitis were examined. Of the total conjunctival scraping samples, 38 of them stained positive with DIF and 29 of them stained positive with Giemsa. The DIF method consumed five more minutes of time compared to Giemsa staining (45 minutes vs. 40 minutes). On the other hand, detecting a positive result in DIF was less difficult compared to detecting one in Giemsa method [13]. In a completely accidental manner (3% of all cases), simultaneous incidence of chlamydia and adenovirus infection is evident and this finding should be anticipated in patients with long-standing follicular keratoconjunctivitis [23]. In conclusion, the Giemsa staining technique is a method with low sensitivity (38.53%)
and high specificity (97.16%) in comparison to DIF. Therefore, Giemsa method is not suitable for routine diagnostic means considering chlamydial conjunctivitis and the possibility of not diagnosing numerous patients infected with C. trachomatis is increased when using this assay. Regarding its ease of performance and high specificity and high sensitivity, DIF is a suitable technique and the method of choice for diagnosis of chlamydial conjunctivitis. However, for a more accurate and more efficient assessment of the sensitivity and specificity of this technique, a parallel study comparing DIF with techniques such as culture and PCR is recommended.

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Footnotes

Authors’ Contribution: Fahimeh Asadi-Amoli and Zohreh Nozarian: Obtaining samples, observation of pathology slides, interpretation of results, drafting the manuscript. Vahid Mehrtash: Translation and language editing. Hooshang Beheshtnejad: Clinical diagnosis and lab patient referral. Avishan Shabani: Statistical calculations.

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References